	1. DISTRICT ADDRESS & PHONE NUMBER					
DEPARTMENT OF HEALTH AND HUMAI FOOD AND DRUG ADMINISTRAT	850 Third Avenue Brooklyn, NY 11232 718-340-7000					
2. NAME AND TITLE OF INDIVIDUAL			3. DATE		4. SAMPLE NUMBER	
Richard A. Frost, General Manager			12-4	-04	25563	
		6. FIRM'S DEA NUMBER				
Quality Wholesale Drug Co.						
7. NUMBER AND STREET	8. CITY AND STATE (Include Zip Code)					
3146 Front Street		Brooklyn, N				
9. SAMPLE COLLECTED (Describe fully. List lot, serial, model Administration and receipt is hereby acknowledged pursuar Federal Food, Drug, and Cosmetic Act [21 USC 360ii(b)] are (NOTE: If you bill FDA for the cost of the Sample(s) listed by	nt to Section 704(c) of nd/or 21 Code of Fede relow, please attach a	the Federal Food, eral Regulations (C copy of this form to	Drug, and Cosmet FR) 1307.02. Exce o your bill.)	ic Act [21 U.S.C. 3 rpts of these are q	374(c)] and / or Section 532 (uoted on the reverse of this f	(b) of the form.
One Box of 25 - 1 cc ampules, Di Knoll Pharmaceutical Co., Orange NJ.		ydromorphi	ne) 2 mg/cc,	lot # 01032	213 manufactured t	эу
	T RECEIVED FOR SA	providing comple to FDA at no aborgo				son
☐ PROVIDED AT NO CHARGE	☑ CASH	BILLED	providing sample	FIOT DA ALTIOUNA	ry <i>e.)</i>	
☑ PURCHASED	□ VOUCH	IER □ CREDIT CARD	Richard	A. Frost	.	
BORROWED (To be returned) \$15.00						
13. COLLECTOR'S NAME (Print or Type) 14. COLLECTOR'S		TLE (Print or Type)	15. COLLECTOR	'S SIGNATURE	
Sylvia A. Rogers		Investigator		Sulvia A Rossus		

Section 704 (c) of the Federal Food, Drug, and Cosmetic Act [21 U.S.C. 374(c)] is quoted below:

"If the officer or employee making any such inspection of a factory, warehouse, or other establishment has obtained any sample in the course of the inspection, upon completion of the inspection and prior to leaving the premises he shall give to the owner, operator, or agent in charge a receipt describing the samples obtained."

Section 532(b) of The Federal Food, Drug and Cosmetic Act [21 USC 360 ii (b)] is quoted in part below:

"Section 532(b) In carrying out the purposes of subsection (a), the Secretary is authorized to-

- (2) ****
- (3) ****
- (4) procure (by negotiation or otherwise) electronic products for research and testing purposes, and sell or otherwise dispose of such products"

21 Code of Federal Regulations 1307.02 is quoted below:

"1307.02 Application of State law and other Federal law. Nothing in this chapter shall be construed as authorizing or permitting any person to do any act which such person is not authorized or permitted to do under other Federal laws or obligations under international treaties, conventions or protocols, or under the law of the State in which he/she desires to do such an act nor shall compliance with such be construed as compliance with other Federal or State laws unless expressly provided in such other laws."

Therefore, in the event any samples of controlled drugs are collected by FDA representatives in the enforcement of the Federal Food, Drug, and Cosmetic Act, the FDA representative shall issue a receipt for such samples on FDA form FDA 484, RECEIPT FOR SAMPLES, to the owner, operator, or agent in charge of the premises.

Report of analysis will be furnished only where samples meet the requirements of Section 704(d) of the Federal Food, Drug, and Cosmetic Act [21 U.S.C. 374(d)] which is quoted below:

"Whenever in the course of any such inspection of a factory or other establishment where food is manufactured, processed, or packed, the officer or employee making the inspection obtains a sample of any such food, and an analysis is made of such sample for the purpose of ascertaining whether such food consists in whole or in part of any filthy, putrid, or decomposed substance, or is otherwise unfit for food, a copy of the results of such analysis shall be furnished promptly to the owner, operator, or agent in charge."